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EP 1 595 908 A1

(12)

EUROPEAN PATENT APPLICATION

(43) Date of publication: 16.11.2005 Bulletin 2005/46 (51) Int CI.7: **C08G 73/00**, C08G 73/02, C08G 73/06, H01B 1/12

(21) Application number: 05010076.7

(22) Date of filing: 09.05.2005

(84) Designated Contracting States:
AT BE BG CH CY CZ DE DK EE ES FI FR GB GR
HUIE IS IT LI LT LU MC NL PL PT RO SE SI SK TR
Designated Extension States:
AL BA HR LV MK YU

(30) Priority: 11.05.2004 KR 2004033168 19.04.2005 KR 2005032461

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(54) Conductive polymers having highly enhanced solubility in organic solvent and synthesizing process thereof

(67) The present invention relates to a new process of synthesizing conductive polymers from monomers substituted with amine group. The process provides simple synthesizing steps for the conductive polymers without using other additives such as stabilizers or emulsifiers. The conductive polymers synthesized according to the present invention have highly enhanced solubility in common organic solvents and electrical conductivity compared to conventional conductive polymers. Therefore, the conductive polymers synthesized according to the present process can be utilized in applications that require high electrical conductivity, for example an electro-magnatic interference shirted or a transparent electrode of thin film, as well as in specific applications such as various conductive films, fibers, polymer blends, battery electrodes or conductive beth mask layers.

$$= \underbrace{ \left(\frac{5}{1} + \frac{2}{3} + \frac{6}{3} + \frac{7}{3} + \frac{3}{3} + \frac{3}{$$

benzenoid ring (phenylene diamine) quinoid ring (quinone diimine)

FIG. 1

Description

[0001] This application claims the benefits of Korean Patent Application No. 2004-33168, filed on May 11, 2004 in Korea and Korean Patent Application No. 2005-0032461, filed on April 19, 2005, which are herein incorporated by reference for all purposes as if fully set forth herein.

BACKGROUND OF THE INVENTION

Field of the Invention

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[0002] The present invention relates to conductive polymers, and more particularly to conductive polymers which have highly enhanced solubility in organic solvents and electrical conductivity, and synthesizing process thereof.

Discussion of the Related Art

[0003] Conductive polymers have conjugated structures along double bonds present in the backbone thereof and has much enhanced electrical conductive properties compared to other organic materiate because the conductive polymers form partial electrical charges along the conjugate structures and thereby having unlocalized electrons when the polymers are doped with dopents such as a protonic acid. Because the conductive polymers have both enhanced electrical, magnetic or optical properties comparable with conventional metals and satisfactory mechanical properties and processability as conventional polymers, they have been remarkably attracted in the filed of chemistry, physics, material enoineering and industries.

[0004] The first developed conductive polymer is polyacetylene, which was developed by Shirakawa et al., however, polyacetylene is oxidized easily in the air. After polyacetylene was developed, Conductive polymers such as polyaniline, polypyrrole, and polythiophene have been developed.

[0005] The conductive polymers can be used in various applications according to their electrical conductivity. For example, the conductive polymers with electrical conductivity of $10^{-13} - 10^{-7}$ (Sicm), $10^{-6} - 10^{-2}$ (Sicm), and equal to and more than 10° (Sicm), respectively have been used as antistatic materials, static discharge materials, and electromagnetic interference(EMI) shielding materials, battery electrodes, semiconductor and solar cells. Accordingly, the conductive polymers may be utilized in more various applications by improving their electrical conductivities.

[0006] Among intrinsically conducing polymers, polyaniline has been noticed in the relevant field since it is not only cheap and very stable compared to polypyrrole and polythiophene also doped easily by protonic acids.

[0007] The polyaniline (PANI) can be classified into the completely reduced from, leucoemeraldine, the intermediated oxidized form, emeraldine, and the fully oxidized form, pemigraniline, according to its oxidation state.

[0068] However, the conductive polymers synthesized through the conventional processes, especially the polyaniline as the completely reduced from, leucoemeraldine, the infermediated oxidized from, emeraldine salt, and the fully oxdized form, pemigraniline, have disadvantages that they cannot be made from melting process owing to their high boiling point and that they must experience complex processing steps since they have low solubility in solvents with high-boiling point or universal or commatible solvents such as meta-cressol.

[0009] In order to improve the problems of the conductive polymers as indicated above, copolymers such as aniline derivatives or graft copolymers have been synthesized by inducing various side chains into the benzene ring or amine group of the conductive polymers for improving solubility of the backbone of the conductive polymers for improving the processability and the electrical conductivity of the conductive polymers. However, those composites have lower electrical conductivity compared to the conductive polymers before reforming.

[0010] Polyaniline (PANI) can be synthesized either by electro-chemical charge transfer reaction which uses electrochemical reaction or by chemical oxidation process is alth uses protonation through acid-base or redox reaction. However, it has been known that the chemical oxidation process is suitable for producing onlyaniline in industrial scale.

[0011] Representative chemical oxidation process for synthesizing polyaniline has been reported to MacDiarmid et al., who synthesized polyaniline by polymerizing aniline monomers dissolved in hydrochloric acid with oxidizing agents such as ammonium persuffate in aqueous solution in the temperature of 1 - 5 °C, separating and washing the precipitates and then obtaining polyaniline (See A.G. MacDiarmid, J.C. Chiang, A.F. Richter, N.L.D. Somarisi, in L. Alcacer (ed.), Conducting Polymers, Special Applications, Reidel, Dordercht, 1987, p. 105). The MadDiarmid process have been utilized widely and regarded as a standard method for producing polyaniline.

[0012] The polyamiline of emeraldine base (EB) synthesized according to the MacDiarmid process has low molecular weight (infinized viscosity 0.8 – 1.2 d/lg), but it is dissolved in 1-methyl-2-pyrotion (NMP). Jost it has reported that emeraldine salt produced by doping the EB with 10-camphorsulfonic acid (ES-CSA) is dissolved at little in meta-cresol. The film made from that solution containing ES-CSA has at most electrical conductivity of about 100 S/cm, on the other

hand, the film made from emeraldine salt doped with hydrochloric acid (ES-HCI) shows highly lower electrical conductivity of about 5 S/cm. However, it needs to be separating not dissolving portion from the dissolved portion in the MacDiarmid process. Especially, the polyaniline synthesized according to the MacDiarmid process has low molecular weight, broad molecular weight distribution, and inferior solubility to solvents or electrical conductivity resulted from side chain reactions to the backbone. Therefore, there remains a need of improving the micro-chemical structure or electrical conductivity of polyaniline synthesized according to the MacDiarmid process.

10013] In order to improve the disadvantages and infenior processability of the polyaniline synthesized by MacDiarmid process, a lot of researches which use emulsion polymerization have been suggested. For example, U.S. patents No. 5,224,631 and No. 5,324,463 to Cao et al., which are incorporated herein by reference, disclose process for synthesizing polyaniline by dissolving aniline monomers and functionalized protonic acid in polar solvents such as water mixing the solution with an organic solvent to prepare an emulsion, and then adding an oxidizing agent into the emulsion. Cao et al. reported that the emeraldine salt (ES) can be dissolved in nonpolar solvent such as xylene because emulsifier acts as a dopant, and therefore, it is reacted with the polyanitine to form composite. However, since Cao et al. uses functionalized protonic acids as emulsifier, it is difficult to control doping the emulsifier and the process requires commonly expensive material. Further, since the functionalized organic acid is hardly separated from polyamiline after polymerizing reaction, the conductive polymers may have only relimited uses and highly inferior electrical properties. For instance, the emeraldine salt, which synthesized according to Cao et al, doped with oddecyl benzene suffonic acid (DSS) has a solubility of less than 0.5 % and an electrical conductivity of only about 0.1 S/cm.

[0014] Kinlen of Monsanto produced polyaniline salt by preparing reverse emulsion system comprising an organic solvent such as 2-butoxyethanol soluble in water and an organic acid, which is not soluble in water but soluble in the organic solvent, as a hydrophobic emulsifier, mixing an aniline monomer and a radical initiator with the emulsion system and polymerizing the mixture to form polymer solution that has an organic layer, which contains polyaniline salt, separated from an aqueous layer containing the radical initiator and non-reacting compounds. (See U.S. patent No. 5,587, 356, Kinlen, Macromolecules, 31, 1745 (1998), which are incorporated herein by reference). Kinlen reported that the polyaniline salt was soluble in nonpolar solvents of no less than 1% (w/w). However, it is difficult to synthesize polyaniline because the radical initiator in the aqueous layer is separated from the monomer in the organic layer and polyaniline salt synthesized orductivity owing to difficulty of control doping process. For example, it was reported that polyaniline salt synthesized with dinonyl naphthalene sulfonic acid as a hydrophilic organic acid had an electrical conductivity of work in examples.

for [0015] Harfev et al. synthesized polyaniline salt with the MacDiarmid process except using pyruvic acid instead of hydrochloric acid (See U.S., patent No. 5,618,469, which is incorporated herein by reference). It is possible to improve processability of polyaniline by using pyruvic acid because pyruvic acid functions as organic solvent as well as dopant. However, since pyruvic acid has lower acidity it is difficult to dope polyaniline by pyruvic acid. Accordingly, polyaniline doped with pyruvic acid has low electrical conductivity, and especially in case the polyaniline doped with pyruvic acid is used as a transparent electrode, it has very high apparent surface resistance as much as 20,000 \(\Omega / \) square, which is very high electrical resistance for the transparent electrode.

[0016] Ho et al., produced polyaniline through emulsion system prepared by adding specific emulsifier into an organic mixture solvent comprising an aniline monomer and a protonic acid with stirring (See U.S. patent No. 6,030,551, which is incorporated herein by reference). According to Ho et al., both a radical initiator such as benzoyl peroxide and the polyaniline is dissolved in the same non-aqueous layer, and therefore, it is possible to synthesize polyaniline solution in situ without residual solids. However, since it is not easy to separate the non-aqueous layer from an aqueous layer, it is expected that polyaniline synthesized according to Ho process may not have high electrical conductivity.

[0017] U.S. patent No. 6,072,027 to Carey et al., which is incorporated herein by reference, discloses a producing method of polyamiline with highly enhanced polymeration yield, by using chlorate salt or hydrochloric acid combined with bi- or trivalent iron salt as a new oxidization initiator.

[0018] Palaniappan et al. disclose a process for the preparation of polyaniline salt by forming inverted emulsion system that comprises an aqueous layer and an organic layer using a surfactant and then polymerizing the inverted emulsion system at room temperature using a radical initiator such as benzoy peroxide dissolved in the organic layer (See U.S. patent publication No. 2002-00062005, U.S. Patents NO. 6,586,565 and 6,50,567, which are incorporated herein by reference). However, polyaniline film repeared from Palaniappan process has very low electrical conductivity, for example about 0.1 \$/cm, and may be only used in much limited applications since it is impossible to raise molecular weight of polyaniline.

[0019] In addition to emulsion polymerization as above, polyaniline synthesis processes through a dispersion polymerization, in which monomers such as aniline is fully dissolved in reacting solvent while synthesized polymers are not dissolved in the solvent, have been reported. For example, Armes et al. reported the polymerization process which comprises stabilizing sterically the conductive polymer by designing particular stabilizer and then particularizing the conductive polymer (See Armes et al., handbook of Conducting Polymers, Elsenbaumer ed. M. Dekker, New York, 1995, Vol. 1, p. 423). In this dispersion polymerization, since most of the stabilizer covers with the polyaniline, the

polyaniline in aqueous solution can be prepared. However, the synthesized polyaniline has a particle size of about 60~300 nm, which is affected by the stabilizer, and has low electrical conductivity, which defines its application.

[0020] Further, there have been reported that polyaniline is synthesized in aqueous solution containing organic solvents. Geng et al., prepared polyaniline film, which has electrical conductivity of about 10 S/cm, through synthesizing polyaniline with organic solvents such as eithanol, THF, and acetone (See Geng et al., Synth. Metals. 86, 1 (1989). However, since it needs very long polymerization reaction time in Geng process, a probability of side reaction is resided. [0021] According to Beadel et al., the polyaniline produced by the standard synthesizing method disclosed in MacDiarmid as described above has higher electrical conductivity as it has higher molecular weight. Accordingly, the monomer needs to be reacted or polymerized at lower temperature in order to enhance molecular weight of the polymerize (See Beadel et al., Synth. Met. 95, 29 – 45, 1998). For lowering reacting temperature, when aniline monomer is polymerized in homogeneous aqueous solution system, metallic salts such as LIC, CaF₂ and the likes are usually added to the system in order to prevent the system from freezing. However, mixing those metallic salts with the solution system causes the reaction to being skow, that is to say, at least 48 hours to complete the polymerizagi reaction, and therefore, it is difficult to control the polymerizagi root proveniline has an increased molecular weight as well as molecular weight distribution (polydispersity of equal to or more

[0022] Also, there form side chains as the aniline monomer is added into a quinonedimine group in intermediate chains. Accordingly, FeCl₂ as an oxidizing agent is added during polymerization reaction in order to inhibit formations of the side chains in polyaniline, or the polyaniline is eluted with organic solvents for removing side products such as oligomers which quit synthesis during the polymerization reaction. Besides, since the monomers are added into the polyaniline the orthor-positions as much as the parapositions of the benzaen ring in the polyaniline backonic as a much as the parapositions of the benzaen ring in the polyaniline has much side chains, which cause the polyaniline to have lower electrical conductivity and solubility.

than 2.5).

Huang et al., J. Am. Soc. 125, 314 (2003)).

[0023] According to Thyssen et al., there is a probability of about 10% of the ortho coupling, which induces side chains in the backbone of the polymers, when the aniline monomers are polymerized by using electro-chemical prosess (See Thyssen et al., Synth. Met. 29, E357 – E362, 1989). Such polymers synthesized by ortho-coupling has lower hydraulic dimensions, which results in decreased intrinsic viscosity, compared to polymers synthesized by para-coupling, i.e. polymers without side chains. In other words, the polymers synthesized by ortho-coupling has much side chains and has more molecular weights even though they have low intrinsic viscosity of equal to or less than 1.2 dlig. Accordingly, the polymers synthesized by ortho-coupling has inferior processability without improving the electron-coupling has inferior processability without improving the electron-

conductivity.

[0024] Moreover, Huang et al. produced polyaniline of nano-fiber form by preparing a system which comprises an organic layer and an aqueous layer immiscible with the organic layer, dissolving an aniline monomer in the organic layer, and polymerizing the monomer in the interface solven.

[0025] Min of Dupont Technology reported that conductive polymer with high yield could be obtained by increasing level of LiCl or NaCl as additive in the MadDlarmid process, as described above, up to 5-10 M at 0 °C for 3 hours (See G. Min. Synth, Met. 119, 273, (2001)).

[0026] In addition to the patents and references described above, many researches were reported for improving physical or chemical properties, for example electrical conductivity of the conductive polymers (Organic Conductive molecules and Polymers, Vol. I - IV, Ed. By H. S. Nalwa, John Wiley & Sons, New York, 1997; Handbook of Conducting Polymers Vol. I, II, Ed. By Skotheim et al., Marcel Dekker, New York, 1998; Conductive Polymers, P. Chandrssekhar, Kluwer Acade. Pub. Boston, 1999; Conductive Electroactive Polymers by G.G. Wallace, G.M. Spinks, L.A.P. Kane-Maquire, P.T. Fasadale, 2nd ed. CRC Press, New York, 2003.

5 [0027] However, the polyaniline producing processes disclosed to date make use of introducing substituent into monomers or mixing the monomers with immense amount of additives such a stabilizer or emulsifier, and therefore, it is difficult to obtain pure polyaniline. Also, because polyaniline according to conventional processes has synthesized by ortho-coupling as much as para- coupling and frequently forms side chains by side reactions, such polyaniline does not have high electrical conductivity, which limits its applications.

[0028]. Also, polypyrole has been synthesized mainly by electro-chemical synthetic process. In case of synthesizing polypyrrole by the electro-chemical process, unlike synthesizing polyparline, acids are not added during polymerization, which makes the reaction simplify. However, when polypyrrole is synthesized according to the chemical process, side reactions such as inter-chain crosslink or side chain addition to the backbone of polypyrrole frequently happened, which causes synthesized polypyrrole to yel dissolving in common solvents and therefore deterrolates processability, in case of synthesizing polypyrrole by electrical process, solvents or counter ions of conductive plate have highly affects on physical properties of polypyrrole.

[0029] Lee et al. prepared of conductive polypyrrole powder by reacting pyrrole monomers at 0 °C for 40 hours using chloroform and dodecyl benzene sulfonic acid (DBSA) of the same molar equivalents of chloroform (See J.Y. Lee, D.

Y. Kim, C.Y. Kim, Synth. Met. 74, 103 (1995)). DBSA adopted by Lee et al. acts as both dopant and surfactant. However, polypyrrole film sample synthesized by Lee et al. has very low electrical conductivity of about 5 S/cm.

[0030] Besides, chemical process for synthesizing polypyrrole in organic solvents such as CHCl₃, THF, or CH₃N₂O has been tried in order to produce polypyrrole being able to dissolve in the organic solvents. However, such synthesized polypyrrole did not have electrical conductivity at all.

[0031] Ames et al. reported a process for preparing stable colloidal polypyrrole by using poly-vinylalcohol, polyetihyleneoxide, or poly-vinylpyridine as a steric stabilizer dissolved in water (See Armes et al., Handbook of Conducting Polymers Elsenbaumer ed. M. Dekker, New York, 1996, Vol. 1, p. 423). However, since polypyrrole powders are surrounded by a lot of stabilizers, like polyaniline, and therefore polypyrrole has a very low electrical conductivity.

[0032] Accordingly, it may enhance electrical conductivity of polypyrrole by linking pyrrole monomers on 2, Spositions between pyrrole ring and sustaining linearity thereof. As mentioned above, pyrrole may be soluble in much solvents compared to aniline, however, it is very difficult to dissolve oxidizing agent and pyrrole monomer in the same solvent. [0033] The synthetic conductive polymers, especially polyaniline has much lower real electrical conductivity than theoretically calculated electrical conductivity, about 105 ~ 105 Scm (Kohlamu et al., Phys. Rev. Lett. 78(20, 3915, 1997), because they does not have fully linear form and form completely order as uch as crystaline structure per se. Since such polymers with lower electrical conductivity cannot be utilized as transparent plastic electrode or EMI shield-ring materials, there still remain needs of development of polymeriline having much improved electrical conductivity in

SUMMARY OF THE INVENTION

the related field.

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[0034] Accordingly, the present invention is directed to conductive polymers with much enhanced electrical conductivity and solubility to common solvents and process for producing the polymers that substantially obviates one or more of problems due to limitations and disadvantages of the related art.

[0035] The present invention is based on a new concept of self-stabilized dispersion polymerization (referred to as "SSDP"). The use of the term "self-stabilized 'herein includes, but is not limited to, the dispersion in the absence of any stabilizers. For example, in contrast with conventional homogeneous or dispersion polymerization using an aqueous medium containing aniline, pyrrole, acid, and oxidant, this new polymerization process is performed in a heterogeneous biphasic system of organic and aqueous medium without any stabilizing additives. Here, the monomer and growing polymer chains act as a stabilizer, resulting in excellent dispersion of the organic phase inside of the aqueous reaction medium.

[0036] It is an objective of the present invention is to provide a synthesizing process of conductive polymers without using other additives such as emulsifier and antifreeze thereby reducing polymerization reaction time. Accordingly, the conductive polymers of the present invention have much highly improved physical property, for example electrical conductivity and solubility to common solvents.

[0037] It is another objective of the present invention is to provide a conductive polymer that has highly improved micro-chemical structure with little structural defects.

[0038] Additional features and advantages of the invention will be set forth herein which follows, and in part will be apparent from the description, or may be learned by practice of the invention. These and other advantages of the invention will be realized and attained by the structure particularly pointed out in the written description and claims hereof as well as the appended drawings.

[0039] To achieve these and other advantages and in accordance with the purpose of the present invention, as embodied and broadly described, in one aspect, the present invention provide a process of synthesizing a conductive polymer, the process comprising: (a) mixing a monomer containing an amine group and an organic solvent with an acid solution; and (b) adding a radical initiator dissolved in a protonic acid into the acid solution to synthesize the conductive polymer.

[0040] Preferably, the monomer is mixed with the acid solution prior to the organic solvent.

[0041] Especially, the monomer with or without substituents has a structure represented by formula I below.

$$H \xrightarrow{R_s} R_s$$

$$R_s = R_s$$

wherein R_i is hydrogen, alkyl, or alkoxy group; and each R_2 to R_5 is respectively hydrogen, alkyl, alkenyl, cycloalkyl, cycloalkenyl, alkyl-thioalkyl, alkyl-anyl, alkyl sulfonyl, alkyl sulfonyl, alkyl sulfonyl, alkyl, hydroxyl, halgen, nitro, or alkyl-aryl.

[0042] Also, the monomer with or without substituents has a structure represented by formula II below.

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$$\begin{array}{c|c}
R_3 & R_2 \\
N & R_3
\end{array}$$
(II)

wherein R₁ is hydrogen, alkyl, or alkoxy group; and each R₂ and R₃ is respectively hydrogen, alkyl, alkenyl, cycloalkyl, cycloalk enyl, alkyl-thioalkyl, alkanoyl, thioalkyl, aryl-alkyl, ankly-amino, amino, alkoxy carbonyl, alkyl sulfonyl, alkyl sulfinyl, thioaryl, sulfonyl, carboxyl, hydroxyl, halogen, nitro, or alkyl-aryl.

[0043] It is preferable that the acid used in step (a) of the present invention may comprise inorganic acid, and more preferably, the acid is selected from the group consisting of hydrochloric acid, sulfuric acid, nitric acid, or phosphoric acid.

[0044] Also, the protonic acid in step (b) of the present invention may be an inorganic acid such as hydrochloric acid, nitric acid, sulfuric acid, phosphoric acid, hydroflycolic acid, or hydroidic acid. The protonic acid of the present invention comprises an organic acid, and preferably, the organic acid is selected from the group consisting of methyl sulfonic acid, dodecyl benzene sulfonic acid, antraquinone-2-sulfonic acid, 4-sulfosalicylic acid, camphor sulfonic acid, chlorinated sulfonic acid, triflucro-sulfonic acid.

[0045] It is characterized that the organic solvent in step (a) has a solubility factor of about 17 to about 29. The organic solvent comprises hydrocarbons unsubstituted or substituted with hydroxyl, halogen, oxygen, ketone, or carboxyl group, such as an alkyl halide.

[0046] The hydrocarbons substituted with halogen may comprise dichloromethane, pentachloro ethane, 1,1,2,2-telrachloro ethane, trichloro ethane, trichloro ethylene, dichloro methane, chloroform, ethyl bromide, ethyl chloride, dichloro propane, trichloro ethane, bis(2-chloroethylether, dichloro ethyl ether, 1,2-dichloro benzene, or mixtures thereof.

[0047] The organic solvent substituted with hydroxyl group may be selected from the group consisting of comprise 1-propanol, 2-methyl-2-propanol, 1,2-dipropandiol, 1,3-propandiol, isopropyl alcohol, butanol, neopentanol, 2-methya-ethanol, 2-butoxy ethanol, 2-ethyl-1-butanol, 3-methyl-1-butanol, 2-methya-2-butanol, 3-methyl-2-butanol, 1-pentanol, 2-pentanol, 3-pentanol, 1,2-propanediol, 1,5-pentandiol, amylalcohol, 2-methyl-1-pentanol, 3-methyl-1-pentanol, 3-methyl-2-pentanol, 3-methyl-3-pentanol, 3-methyl-3-

[0048] The organic solvent substituted with oxygen may comprise ethylene glycol monoethyl ether, ethylene glycol dimethyl ether, ethylene glycol monomethyl ether, ethylene glycol monoethyl ether, diethylene glycol monobutyl ether, diethylene glycol diethylene glycol dimethyl ether, diethylene glycol diethylene glycol dimethyl ether, diethylene glycol monomethyl ether, diethylene glycol monoethyl ether, diethylene glycol monoe

[0049] The organic solvent substituted with ketone group comprises butly methyl ketone, methyl-ethyl ketone, 4-hydroxy-4-methyl-2-pentanone, cydopentanone, diacetone alcohol, 4-methyli-pentanone, 4-methyl-2-pentanone, or mixtures thereof.

[0050] Also the organic solvent may comprises common organic solvents such as dietrly carbonate, benzyl acetate, dimethy glutarate, ethylacetoacetate, isobutyl isobutanoate, isobutyl acetate, meta-cresol, toluene, xylene, nitrobenzene, tetrahydrofuran, N-methyl-2-pyrolidone, dimethyl sulfoxide, N,N-dimethylformamide, or mixtures thereof.

[0051] Moreover, the radical initiator comprises ammonium persulfale, hydrogen peroxide, manganese dioxide, pottassium dichromate, potassium choride, potassium permanganate, potassium bromate, potassiu

[0052] It is also preferable that step (b) of the present invention is performed in the temperature of between about -45 °C to about 40 °C. Also, the radical initiator and the organic solvent comprises an organic phase, wherein the

organic phase comprises about 5~95 % by weight based upon total aqueous solution.

[0053] It is more preferable that the process of the present invention further comprises step (c) dedoping the conductive polymer with a base such as hydroxide compounds.

[0054] In another aspect of the present invention provide a conductive polymer synthesized by the present process, wherein the conductive polymer has a hollow quadra-angular rod shape and honeycombed network configuration. The conductive polymer synthesized by the present invention is consisted of nanometer particles, and has an apparent density in the range of about 0.03 ~ 0.19 measured in ASTM Standard D1895-6.

[0055] In still another aspect, the present invention provides a electrical conductive polymer synthesized according to the process above, wherein the polymer has an electrical conductivity of at least about 300 S/cm. Preferably, the conductive polymer has an electrical conductivity of at least about 500 S/cm, for example at least about 700 S/cm or at least 900 S/cm, more preferably, at least about 1100 S/cm, and most preferably, at least about 1300 S/cm.

[0056] In further another aspect, the present invention provides a conductive polymer synthesized according to the present invention, wherein the conductive polymer has a hollow quadra-angular rod shape and honeycombed network configuration, wherein the conductive polymer has a repeat unit represented by the formula III below and the conductive polymer has at least one single peak at about 123 ppm of chemical shift and at about 158 ppm of chemical shift in a ¹³C CPMAS MMR spectrum and/or has identifiable peaks at around 140 ppm of chemical shift in a ¹³C CPMAS MMR spectrum and/or has identifiable peaks at around 140 ppm of chemical shift in a 13C CPMAS MMR spectrum.

wherein x and y is respectively a molar fraction of quinonediimine structural unit and phenylenediamine structural unit in the repeating unit, and 0 < x < 1, 0 < y < 1 and x + y = 1; and n is an integer of 2 or more.

[0057] The conductive polymer forms peaks at about 138 ppm of chemical shift and at about 143 ppm of chemical shift in a ¹³C CPMAS NMR spectrum. Particularly, the conductive polymer has 1₁₃₈ larger than 1₄₅₂ wherein 1₁₃₈ represents a peak intensity at about 138 ppm of chemical shift in the ¹³C CPMAS NMR spectrum and 1₄₅₂ represents a peak intensity at about 143 ppm of chemical shift in the ¹³C CPMAS NMR spectrum. Preferably, the conductive polymer has a peak intensity ratio, 1₁₃₈ / 1₁₄₃, of equal to or more than 1.2 in the ¹³C CPMAS NMR spectrum. Besides, the conductive polymer has to peaks at about 1107 cm⁻¹ of weakength in PAS spectrum.

[0058] In further still another aspect, the present invention provides a polyaniline having a repeat unit represented by the formula below, wherein the polyaniline has three main peaks corresponding to quaternary carbon in a solution state ¹⁹C NMR spectrum in case the polyaniline is substituted with terl-butoxycarbonyl.

[Formula]

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wherein x and y is respectively a molar fraction of a quinonediimine structural unit and phenylenediamine structural unit in the repeating unit, and 0 < x < 1, 0 < y < 1 and x + y = 1; and n is an integer of 2 or more.

[0059] The conductive polymer synthesized according to the present invention has highly linear configuration, fewer side chains, and therefore highly improved electrical conductivity compared to polyaniline synthesized according to conventional process. Accordingly, polyaniline of the present invention may be used as various conductive films, fibers, coatings, blends with other polymers, battery electrodes, or material for organic semiconductors or organic descolar Especiality colvaniline synthesized according to the present invention may be utilized as transparent electrodes soice.

cells, conductive etch mask layer or for anti-corrosion, absorbency of near infrared light since composites or composition comprising polyaniline of the present invention has highly improved electrical conductivity even though low contents of polyaniline.

[0060] It is to be understood that both the foregoing general description and the following detailed description are exemplary and explanatory and are intended to provide further explanation of the invention as claimed.

BRIEF DESCRIPTION OF THE DRAWING

[0061] The accompanying drawings, which are included to provide a further understanding of the invention and are incorporated in and constitute a part of this specification, illustrate embodiments of the invention and together with the description serve to explain the principles of the invention.

[0062] In the drawings:

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[0063] FIG. 1 is a schematic chemical structure showing the repeating unit of polyaniline synthesized according to a preferred example of the present invention for describing its particular chemical micro-structure;

[0064] FIG. 2 shows a spectrum resulted from ¹³C CPMAS NMR analysis for highly conductive polyaniline (HCPANI) synthesized according to a preferred example of the present invention;

[0065] FIG. 3 shows a spectrum resulted from ¹³C CPMAS NMR analysis for polyaniline (PANI) synthesized according to a conventional process;

[0066] FIG. 4 shows a spectrum resulted from PAS analysis for highly conductive polyaniline (HCPANI) synthesized according to a preferred example of the present invention:

[0067] FIG. 5 shows a spectrum resulted from PAS analysis for polyaniline (PANI) synthesized according to a conventional process:

[0068] FIGS. 6A to 6E show respectively SEM electron microscopy of highly conductive polyaniline of emeraldine base form (HCPANI) synthesized according to the preferred examples of the present invention;

[0069] FIGS. 7A to 7E show respectively SEM electron microscopy of polyaniline of emeraldine base form(PANI) synthesized according to the conventional process;

[0070] FIG. 8 shows a spectrum resulted from UV-VIS-NIR analysis for emeraldine base doped with camphor sulfonic acid (EB-CSA) synthesized according to the present invention;

[0071] FIG. 9 a spectrum resulted from solution state ¹³C NMR analysis for polyaniline synthesized according to the preferred example of the present invention; and

[0072] FIG. 10 is a graph showing the molecular-weight distribution of highly conductive polyaniline (HCPANI) synthesized according to the present invention.

DETAILED DESCRIPTION OF THE ILLUSTRATED EMBODIMENTS

[0073] Prior to describing the present invention, polyamiline or its derivatives synthesized according to the present invention will be frequently referred to as "FCANIT," while the polyamiline synthesized according to the conventional method will be referred to as "FANIT" without indicated otherwise herein. Also, polypyrrole or its derivatives of the present invention will be frequently referred to as "FPy" without indicated otherwise herein. In other words, 'HCPANIT' is intended herein to mean conductive polymer synthesized from aniline monomer within is unsubstituted or substituted as mentioned formula I below, and refer to each or all of leucoemeraldine form, emeraldine base (EB), emeraldine salt (ES) or pemigraniline form according to the present invention.

wherein R, is hydrogen, alkyl, or alkozy group; and each R₂ to R₃ is respectively hydrogen, alkyl, alkenyl, cycloalkyl, cycloalkenyl, alkyl-thioalkyl, alkanoyl, thioalkyl, aryl-alkyl, alkyl-amino, amino, alkozy carbonyl, alkyl sulfonyl, alkyl sulfinyl, thioaryl, sulfonyl, carboxyl, hydroxyl, halogen, nitro, or alkyl-aryl.

[0074] Also, 'PPy' is intended herein to mean conductive polymer synthesized from anline monomer which is unsub-

stituted or substituted as described in formula II below according to the present invention.

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wherein each R₁ to R₂ is respectively as defined in formula I above.

[0075] In the process according to the preferred embodiment of the present invention, reactants containing monomer are mixed with 2-phase reaction system that comprises an aqueous phase and an organic solution phase. Accordingly, the present process affers from the MacDiamnid standard polymerizing process. Also, other additives, for example emislifier, polymeric stabilizer, monomeric and/or oligomeric stabilizer, or other templates, are not required principally in the synthesizing process of the present invention. Therefore, the present process is different substantially from the conventional emulsion polymerization, suspension polymerization, or dispersion polymerization as described above. This new polymerization based upon self-stabilized dispersion concept is referred to as SSDP. The use of the term self-stabilization includes, but is not limited to, the stabilization of biphasic reaction systems by the reactants and polymerization products i.e., the absence of any stabilizers or antifreezes or templates. It is possible that both of two phases of the SSDP process consist of non-continuous phases or templates. It is possible that both of two phases of the SSDP process consist of non-continuous phase, however, the aqueous phase may be continuous phase and the organic solution phase may be non-continuous phase, however, the aqueous phase may be continuous phase and the organic solution phase may be non-continuous phase, and vice versa. Particularly, the organic solution phase may be another organic solution of the present invention comprises a hydrophilic solvent such as water, a monomer, for example a monomer defined in formula I or formula II above, and an acid, preferably protonic acid, in initiating reaction.

00771 The hydrophilic solvent may comprise water, methanol, ethanol, acetonitrile, 2-methoxy ethanol, or mixtures thereof, and preferably water atone. The acid may be an inorganic acid or an organic acid which preferably has p.Kx. of equal to less than 4.0, more preferably equal to or less than 3.5, and most preferably a protonic acid. Particularly, the acid may comprise an inorganic acid selected from the group consisting of hydrochloric acid, suffuric acid, nitric acid, or phosphoric acid, or an organic acid such as an any sulforicia acid or any lafty sulforion acid, which is unsubstitude or substituted with halogen, or mixtures thereof. More specifically, the organic acid may comprise an alky sulforic acid such as methyl sulforic acid, ethanol sulforic acid, altoparated alkyl sulforic acid, for example chloro sulforic acid, surfuse the sulforic acid, ethanol such as dodecyl benzene sulforic acid, antirraquinone2-sulforic acid, 5-sulfossilcylic acid or camphor sulforic acid, or mixtures thereof. Preferably, the acid may comprise an inorganic acid such as hydrochloric acid, sulforic acid, intric acid, or mixtures thereof. Preferably, the acid may comprise an inorganic acid such as hydrochloric acid, sulforic acid, intric acid, or or phosphoric acid.

[0078] The organic solution phase in the reaction system of the present invention comprises an organic solvent, preferably immiscible or miscible a little with the aqueous phase or an organic solvent that may be separated and dispersed with the aqueous phase by soluble self oriented material, which is disclosed in international patent publication NO. WO-02/074833 to the present inventors.

[0079] In case of choosing an organic solvent that may be used in synthesizing polymers, a solubility parameter, which is related with the g molecular weight, density of the polymers, should be considered. Reactions using various organic solvents also produce polyanilines of improved chemical micro-structure with similar shapes and conductivity. In a preferred embodiment of the present invention, the organic acid constituted with the organic solution phase may comprise an organic solvent having the solubility parameter of between about 17 and about 29.

[0080] In a preferred embodiment of the present invention, the organic solvent may comprise hydrocarbons, such as aliphatic, alicyclic, or aromatic hydrocarbons, unsubstituted or substituted with hydroxyl, halogen, oxygen, extens, or carboxyl group, or common organic solvents that may be used in synthesizing conductive polymers. The hydrocarbons may be halogen-substituted hydrocarbons such as alkyl halides, ether, alicyclic hydrocarbons or aromatic hydrocarbons. The hydrocarbons or promises C₂-C₃-C₃ clohofols.

[0081] Preferably, the halogen-substituted hydrocarbons comprise a) alkyl halides such as dichloromethane, pentachloro ethane, 1,1,2,2-tetrachloro ethane, trichloro ethane, trichloro ethylene, dichloro methane, chloroform, ethyl bromide, ethyl chloride, dichloro propane, trichloro ethane, or mixtures thereof; b) ether such as bis[2-chloroethyl)ether, dichloro ethyl ether, or mixture thereof; and c) aromatic hydrocarbons such as 1,2-dichloro benzene. Besides, the organic solvent substituted with hydroxyl group may comprise 1-propanol, 2-methyl-2-propandio, 1,2-dipropandiol,

1,3-propandiol, isopropyl alcohol, butanol, neopentanol, 2-methoxy ethanol, 2-butoxy ethanol, 2-ethyl-1-butanol, 3-methyl-1-butanol, 3-methyl-2-butanol, 3-methyl-2-butanol, 1,2-pentanol, 2-pentanol, 3-pentanol, 1,5-pentanolol, 3-methyl-1-pentanol, 2-methyl-2-pentanol, 3-methyl-2-pentanol, 4-methyl-2-pentanol, 3-methyl-2-pentanol, 4-methyl-2-pentanol, 2-methyl-3-pentanol, 3-methyl-3-pentanol, 4-methyl-2-pentanol, 3-methyl-3-pentanol, 3-methyl-3-pentanol, 4-methyl-2-pentanol, 3-hexandiol, adanol, 1-octanol, 2-octanol, decanol, dodecanol, cyclohexanol, tri-ethyl-ene dycol, tetra-tethylene dyco

[0082] Also, the hydrocarbons substituted with oxygen may comprise ethylene glycol monoethyl ether, ethylene glycol dimethyl ether, ethylene glycol monoethyl ether, ethylene glycol monobutyl ether, diethylene glycol monobutyl ether, diethylene glycol monobutyl ether, diethylene glycol dimethyl ether, diethylene glycol dimethyl ether, diethylene glycol dimethyl ether, diethylene glycol monoethyl ether, diethylene glycol monoethyl ether, diethylene glycol dimethyl ether, diethylene glycol monoethyl ether, diethylene glycol dimethyl ether, diethylene glycol dimethyl ether, diethylene glycol monoethyl ether, diethylene glycol dimethyl ether, diethylene glycol dimethyl ether, diethylene glycol monoethyl ether, diethylene glycol dimethyl ether, diethylene glycol monoethyl ether, diethylene glycol ether, diethylene glycol monoethyl ether, diethylene glycol m

[0083] The organic solvent substituted with ketone group comprises butyl methyl ketone, methyl-ethyl ketone, 4-hydroxy-4-methyl-2-pentanone, cyclopentanone, diacetone alcohol, 4-methyhl-pentanone, 4-methyl-2-pentanone, or mixtures thereof.

[0884] Besides, the organic solvent can be used in the present invention comprises diethyl carbonate, benzyl acetate, dimethyl glutarate, ethylacetoacetate, isobutly isobutanoate, isobutly acetate, meta-cresol, toluene, xylene, nitrobenzene, tetrahydrofuran, N-methyl-2-pyrolicione, dimethyl sulfoxide, N,N-dimethylformamide, or mixtures thereof.

[0085] Further, the radical initiator, which is added in the aqueous phase according to the present SSDP process, may comprise ammonium peroxisulfate, hydrogen peroxide, manganese dioxide, potassium dichromate, potassium permanganate, potassium bromate, potassium chlorate, or mixture thereof, and preferably ammonium peroxisulfate. In case of using ammonium peroxisulfate as oxidizing agent or radical initiator, two electrons per 1 mole are related, and therefore, radical initiator can be used about 0.1–5 molar equivalent, preferably about 0.1–0.75 molar equivalent (or 1 mole of mnoment).

[0066] Angelopulos et al, reported that the solubility of the synthesized polymers and stability of polymer solution is reciprocal to the amount of the radical initiator (See Angelopulos et al., Synth. Met. 84, 35, 1997). Accordingly, it is important to control the addition procedure of the initiator as well as the amount of the initiator in the present invention. The addition procedure of the initiator may have an affect on the micro-chemical structures of synthesized polymers because the radical initiator has an affect on the hydrolysis of intermediates synthesized in the polymerization step. [0087] The polymerization reaction of the present invention is exothermal reaction, and therefore, preferably the polymerization step.

reactants are stirred during the reaction. The reaction may be performed in the temperature of -45 °C to 45 °C, such as -45 °C to 40 °C. Preferably, the reaction should be performed at suitable temperature, which can be determined by desired molecular weight, molecular weighd distribution, or electrical conductivity of the synthesized polymers, among the above temperature ranges and keep it during the polymerization reaction because the reaction time and the molecular weigh of the polymers depends on the reaction temperature.

[0088] The above reactants are introduced into a reaction vessel to initiate polymerization. After completing the polymerization reaction, the synthesized polymers may be separated with various methods according to desired formation of final product. For example, highly conductive polyaniline (HCPANI) synthesized according to the preferred embodiment of the present invention is washed with uster or methanol, and recovered to obtain emeraldine salt (ES) prowder. The ES prowder is treated with a base for memeraldine base (ES) which is very soluble in organic solvents. The EB form may be doped with various dopants to reprocess it or processed to dope it for various applications. Or the obtained EB may be manufactured easily to be uconversation for form or vertice priorations from the vertice vector process.

[0089] The SSDP process has main advantage of being able to control molecular weight of produced conductive polymers over the conventional methods. In a preferred embodiment of the present invention, conductive polymers with a molecular weight of between about 10,000 and about 385,000 can be obtained by only changing the reaction conditions such as reaction time, temperature, and the likes. Especially, electrical conductive polymers of the present invention had an intrinsic viscosity of about 0.1 to about 2.9 determined at 30° Ce fler the conductive polymers were dissolved in sulfuric acid with a concentration of 0.1 g/dl. Also, the electrical conductive polymers, especially EB, synthesized according to the present invention has remarkably differentiating micro-chemical structure and highly enhanced electrical conductivity compared to EB synthesized according to the conventional methy compared to EB synthesized according to the conventional methy.

[0090] It has been only known that the polyaniline of EB form synthesized according to the conventional method has a fraction ratio of x to y in formula III above of about 1:1. In other words, the micro-chemical structure of the polyaniline has not been discovered. On the other hands, HCPANI, which has highly enhanced electrical conductivity and is synthesized according to the present SSDP invention, has remarkably differentiating chemical microstructure compared to the conventional PANI. The differences in chemical structure between the HCPANI and PANI will be more described in more detail referring the appended drawings.

[0991] It is only known that the polyaniline of EB form synthesized according to the conventional method has a fraction ratio of x to y in formula above of about 1:1. In other words, the microstructure of the polyaniline has not been fully discovered. On the other hands, HCPANI, which has highly enhanced electrical conductivity and is synthesized according to the present invention, has remarkably characteristic behenical microstructure compared to the convenient.

PANI. The differences in chemical structure between the HCPANI and PANI will be more described in more detail referring the appended drawings.

[0092] FIG. 1 is a schematic formula showing a repeating unit of polyaniline with carbon numbers for describing its micro-chemical structure. FIG. 2 shows a spectrum resulted from ¹³C CPMAS NMR analysis for highly conductive polyaniline (HCPANI) synthesized according to a preferred example of the present invention. FIG. 3 shows a spectrum resulted from ¹³C CPMAS NMR analysis for polyaniline (PANI) synthesized according to a conventional process.

[0093] As shown in FIGS. 2 or 3, HCPANI, which synthesized according to the present invention, has two apparent separated peaks around 140 ppm of chemical shift, that is one peak at about 139 ppm (I₁₃₈) in ¹³C CPMAS (Cross-Polarized Magnetic-Angle-Spinning) NMR spectrum (FIG. 2). On the other hand, PANI has blurring multiple peaks at around 140 ppm in ¹³C CPMAS NMR spectrum (FIG. 3).

[0094] According to Raghunathan et al., the two peaks (I₁₃₈ and I₁₄₃ in FIG. 2) at around 140 ppm of chemical shift in ¹³C CPMAS NMR analysis of the polyaniline of EB form corresponds to protonated carbons connected to hydrogen of quinoid ring in repeating unit of polyaniline of EB form shown in FIG. 1 (Raghunathan et al., Synth. Met. 81, 39 ~ 47, 1996; Yasuda et al., Synth. Met. 61, 239 ~ 245, 1993).

[095] However, it is difficult to certify specific peaks at around 140 ppm in ¹³C CPMAS NMR spectrum of the PANI synthesized according to the conventional method because there are many small peaks at around 140 ppm of chemical shift as shown in FIG. 3. On the other hand, it was determined that the HCPANI synthesized according to the present invention has two or more apparently confirmable peaks, 1443 and 1543, a shoulder at around 140 ppm of chemical shift in ¹³C CPMAS NMR spectrum as shown in FIG. 2. Beaties, it was determined that the HCPANI had higher peak intensity at about 138 ppm of chemical shift than peak intensity at about 143 ppm of chemical shift in ¹³C CPMAS NMR spectrum (1543 ppm of chemical shift in ¹³C CPMAS NMR analysis is one characteristic of HCPANI synthesized according to the present invention, which is remarkably different from peak forms in ¹³C CPMAS NMR spectrum (1743 PCNN synthesized according to the conventional method.

[0996] HCPANI synthesized according to the present invention has two noticeably confirmable peaks at about 140 ppm in ¹3C CPAMS NIMR spectrum, because the quinoid ring (quinonedimine structural unit) in the repeating unit of HCPANI in FIG. 1 is connected through immine bonds and thereby not being able to rotate and having bended form of -N= bonding instead of maintaining linear form. Therefore, 4 carbon atoms (C4) on the quinoid ring shown in FIG. 1 isoe equivalences. Accordingly, we may infer that HCPANI synthesized according to the present invention has nearly theoretically ideal structure of polyanitine as shown formula above. On the other hand, since PANI synthesized according to conventional method has defects in quinoid ring, PANI has different structure form the structure of formula above. Accordingly, it is difficult to certify specific peaks at around 140 ppm of chemical shift in ¹³C CPMAS NMR spectrum of PANI. which has many indistinguishable peaks around 140 ppm.

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[0097] Wei et al. reported that Michael addition reaction of aniline monomer might be happened on the quinoid ring as shown below. Therefore, we think that PANI has other micro-chemical structures than HCPANI.

[0098] Further, HCPANI synthesized according to the present invention has a single or unique peak at about 123 ppm of chemical shift or about 158 ppm of chemical shift in ¹³C CPMAS NMR spectrum as shown in FIG. 2. On the other hand PANI synthesized according to the conventional method shows 2 or more unidentifiable peaks at about 123 ppm of chemical shift ¹³C CPMAS NMR spectrum as shown in FIG. 3

[0099] In relation to the differences of peak formation between HCPANI and PANI in ¹²C CPMAS NMR spectrum, a single peak at about 123 ppm of chemical shift in ¹³C CPMAS NMR spectrum corresponds to carbon atoms C1 and C2 of benzenoid ring (phenylenediamine structural unit), which may be rotated a bit in molecules, of the repeating unit of polyaniline in F1C. I HCPANI had an equivalent unique or single peak at about 123 ppm in ¹³C CPMAS NMR spectrum (FIG. 2), while PANI showed divided peaks, not unique peak, at about 123 ppm in ¹³C CPMAS NMR spectrum (FIG. 3). In other words, it is certified that HCPANI of the present invention has equivalent carbon atoms in benzenoid ring, while PANI does not have equivalent carbon atoms in benzenoid ring.

[0100] Yasuda et al., synthesized polyaniline by adopting Cao et al. (Cao et al., Polymer, 30, 2305, 1989) with using

FeCl₃ instead of ammonium persulfate which is commonly used in conventional chemical oxidation method for producing polyaniline (See Yasuda et al., Synth. Met. 61, 239 – 245, 1993). However, solid state polyaniline synthesized according to Yasuda et al. did not have distinguishable peak, but had only indistinguishable small peaks, at about 138 ppm of chemical shift in ¹³C CPMAS NMR spectrum, and showed the peak intensity at about 138 ppm of chemical shift was lower than the peak intensity at about 143 pom of chemical shift.

[0101]. In other words, conventional PANI has many indistinguishable small peaks at about 138 ppm of chemical shift in 13C CPMAS NMR analysis, and the peak intensity at about 138 ppm is weaker than the peak intensity at about 143 ppm of chemical shift. On the other hand, HCPANI synthesized according to the present invention has two or more notioeably distinguished peaks at around 140 ppm in 13C CMPAS NMR analysis. HCPANI synthesized according to the present invention has few defects at carbons on the quinoid rings of repeating units of polyamiline and antime monomers is bonded through para- coupling in polymerization. Such differences in micro-chemical structures cause HCPANI to ahving much higher electrical conductivity compared to conventional PANI.

[0102] Therefore, it is concluded that HCPANI synthesized according to the preferred embodiment of the present invention has no, or tittle effects in carbons formed on the quinoid ring in the repeat unit thereof and that aniline monomer is synthesized to the para position of another aniline monomer or oligomeric anilines in polymerization, and therefore, HCPANI of the present invention has apparent distinguishable peak at each about 155 ppm, around 140 ppm, or about 123 ppm in "2C CPMAS NMR spectrum. Such differences in micro-chemical structure cause HCPANI of the present invention to have much enhanced electrical conductivity compared to conventional polyaniline (PANI). [0103] HCPANI synthesized according to the present SSDP process has noticeably spectrum in PAS analysis, 1508, 4 and 5 are respectively a graph obtained from Photo Acoustic Spectroscopy (PAS) results which are suitable for obtaining infrared spectrum of polymeric powders such as polyaniline. FIG. 4 shows a spectrum obtained from PAS analysis for HCPANI powder synthesized according to the present invention, while FIG. 5 shows a spectrum obtained from PAS analysis for PANI powder synthesized according to conventional method. It is generally known that morphology of samples has a little affect on absorbency of samples and does not have to do with photometry of samples in PAS analysis. The analytical results shown in FIGS. 4 and 5 were obtained by treating the powder forms of HCPANI and PANI under the same analytical conditions, and then comparing quantitatively the infrared absorbencies.

[0104] Among infrared absorbency peaks of PAS analysis in FIGS. 4 and 5, the peak at about 1107 cm⁻¹ of wavelength is assigned to the ring stretching vibration of the amine (C-N) group in the repeating unit of polyamiline. HCPANI powder has other infrared absorbency peaks at about 1107 cm⁻¹ of wavelength than the PANI powder synthesized according to the conventional method. HCPANI synthesized according to the present invention has two separated peaks at about 1107 cm⁻¹ of wavelength (14_{TO}) is relatively week in PAS analysis. (FIG. 4). On the other hand, PANI synthesized according to the conventional method has a unique peak at about 1107 cm⁻¹ or of wavelength and and 1₁₀₇ is relatively week in (FIG. 5).

[0105] In accordance with the preferred example, the peak formation at about 1107 cm⁻¹ of wavelength in PAS spectrum is closely related to electrical conductivity of polymers and resulted from structural differences between HCPANI and PANI. In other words, HCPANI has two relatively weak peaks at about 1107 cm⁻¹, while PANI has a relatively strong unique peak at about 1107 cm⁻¹ of wavelength in PAS spectra.

[016] Besides, it was certified that molecular weight of the conductive polymer synthesized in the preferred example is closely related to the electrical conductivity. For example, HCPANI synthesized according to the preferred example of the present invention with number average molecular weights of about 10,000 ~ 30,000 has electrical conductivity of about 100 ~ 300 S/cm, while HCPANI with number average molecular weights of about 30,000 ~ 89,000 has electrical conductivity of about 300-1300 S/cm. In other words, electrical conductivity depends on the molecular weight.

[0107] Further, the conductive polymers synthesized according to the preferred examples of the present invention have enhanced solubility compared to the conventional polymers. In case HCPANN of conductive EEE form has a number average molecular weight of 15,000, it has solubility of about 10 % (w/w) in NMP at room temperature, which is about twice as solubility of polyaniline or EEB rwing the same molecular weight, which is about 5 % (w/w) in NMP. Especially, HCPANI with a number average molecular weight of 15,000 - 19,000 (intrinsic viscosity of 1.7 - 2.7 d/lg) has solubility of equal to or more than 3 % (w/w) in NMP, which is much higher than solubility of conventional polyaniline with the same molecular weight, which is less than 2 % (w/w) in MMP.

[0108] The solubility differences between HCPANI of the present invention and the conventional PANI result from the structural differences between them as well as from particle forms or formation in the polymerization. It was certified that the conductive polymers synthesized according to the preferred examples of the present invention have specific particle structure or network configuration because they are polymerized in the reaction system which comprises the autueous phase and the organic solution phase but can be self-stabilized.

[0109] FIGS. 6A to 6F respectively shows SEM electron microscopy of highly conductive polyaniline of emeraldine base form (HCPANI) synthesized according to the preferred examples of the present invention, and FIGS. 7A to 7E respectively shows SEM electron microscopy of polyaniline of emeraldine base form (PANI) synthesized according to the conventional process. As shown in FIGS. 6A to 6F, HCPANI of EB form synthesized according to the present

invention has cross-sections with various lengths from about $10 \, nm$ to about $50 \, \mu m$, particularly in particles of barrel shapes (FIGS. 6A and 6C). More specifically, each of the HCPANI of the present invention has common structure of having internal holes like foamed-shapes or honey-comb shapes or having a plurality of compact hollow quadra-angular rod or quadra-angular bar.

[0110] Specifically, HCPANI has a plurality of globular particles of 20 – 80 nm, which is clustered to form specific network configuration like bunches of grapes as shown in FIG. 6E, which shows 30,000 times enlarged SEM electron microscopy of HCPANI. Such a structure allows HCPANI of the present invention to have much increased surface areas compared to conventional polyaniline. In contrast, it was observed that PANI synthesized according to the conventional method forms particles of precipitates as shown in FIGS, 7A and 7B.

[0111] According to Mandal et al., polyaniline synthesized by dispersion polymerization has common compact structure or configuration even though it has various forms such as needle shape, obtong shape, or spherical shape (See Mandal et al., Langmuir 12, p 1585, 1996). Besides, Huang et al. reported that polyaniline of nano-fiber form can be obtained by dissolving respectively an aniline monomer in an organic solvent and an initiator in water and then synthesizing the mixture solution in the interface (See Huang et al., J. Am. Chem. Soc., 126, p 314, 2003, Angew. Chem. Int. Ed., 43, p5817, 2004). However, HCPANI synthesized according to the preferred examples of the present invention has a sort of a network configuration, which results in increasing surface area of HCPANI particles and thereby enhancing solubility thereof. In relation with the network configuration, it was calculated that HCPANI of the present invention had apparent density of between 0.03 g/ml and 0.19 g/ml measured in ASTM standard D1895-6, which is very low compared to conventional polyaniline.

[0112] As mentioned above, the conductive polymers synthesized according to the preferred examples of the present invention has much enhanced electrical conductivity and solubility compared to conventional conductive polymers. However, in case of increasing the electrical conductivity and solubility of conductive polymers, other additives may be added into the reaction system as templates. For example, precursors of soluble self oriented materials such as a precursor represented by the structure below, disclosed in Example 3 of international patent publication WO 02-074833 to the present inventors may be mixed with the monomers as templates into the reaction system.

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wherein each R is $-(CH_2)_nCH_3$. $-O(CH_2)_nCH_3$. $-O(CH_2CH_2)_nOCH_3$, wherein n is integers between 1 and 24. [0113] The soluble self oriented materials represent by the above structure may be added into the reaction system of about 5-30 % by weight, preferably 5 \sim 25 % by weight, and more preferably 10 \sim 20 % by weight based on the monomers.

[D114] Particularly, the soluble self oriented materials can be mixed with the monomers is a material R is $O(CH_2)_c CH_3$ (in si integers between 1 and 25 yil in the structure above. Preferably, the soluble self oriented materials have side clinic (R) of which terminal end may have substituted with sulfonic acid (SO₂H), carboxylic acid (-COOH), benzane sulfonic acid (-COOH) COOH), acromovement; carboxylic acid (-COOH) (-COOH), acromovement; carboxylic in cid (-COOH) (-COOH), acromovement; carboxylic in cid (-COOH) (-COOH), acromovement; carboxylic in cid (-SH) group.

[0115] The synthesizing method of the soluble self oriented material which may be added in the reaction system of the present invention as template is described in detail in international patent publication no. WO 02-074833.

[0116] Moreover, it was determined that HCPANI synthesized according to the present invention has well-defined micro-chemical structures. FIG. 9 shows a spectrum resulted from ¹³C NMR analysis for polyaniline substituted with left-butoxycarbonyl (t-BCC) of the present invention. The micro-chemical structure was investigated for polyaniline derivative prepared by substituting with tert-butoxycarbonyl (t-BCC) to enhance solubility in common organic NMR solvents such as CDCIs.

[0117] HCPANI-IBOC of the present invention has four main distinguishable peaks between at about 139.5 ppm of chemical shift and at about 160 ppm of chemical shift in solution state ¹³C NMR spectrum. Among four main peaks, three peaks formed at about 140 ppm, about 148 ppm, and about 159 ppm of chemical shift in solution state ¹³C NMR spectrum corresponds to quaternary carbons of HCPANI-IBOC. Therefore, it was certified that HCPANI-IBOC of the present invention forms three main peaks relating to quaternary carbons.

The present invention will be explained in more detail through the following nonlimiting examples. However, the present invention will not be limited to the following examples.

EXAMPLES

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Measurement of Electrical Conductivity

[0118] Electrical conductivity of polymers synthesized in the following examples are measured with commonly used four line probe method at room temperature in the condition of relative humidity of about 50%. Carbon paste was used for preventing the polymers from corroding in case of contacting gold wires. The electrical conductivity of film samples with thickness of about 1 ~ 100 µm (micron) (sample thickness: I, sample width: w) was measured by calculating voltages (V), currents (i), and distances (i) between 2 internal electrodes and 2 external electrodes connected to the samples with Keithlev instruments.

Electrical Conductivity = $(I \cdot i) / (w \cdot t \cdot v)$

5 [0119] Electrical conductivity was calculated by the above equation (S/on or Simen/on). Electrical conductivity was also measured with Van der Pauw method, which uses standard four point probe, in order to certify the homogeneity in electrical conductivity of the samples. The 4 point measurement results were in the range within 5%.

SEM Particle Shape Measurement

[0120] Particle formation, structure of configuration of conductive polymers synthesized in the following examples were analyzed with scanning electron microscope (SEM, model no. XL-30, Philips Co.). SEM photographs particles within very restricted region, and therefore, we observed a lot of microscopy for obtaining representative images.

25 Measurement of Molecular Weight with GPC

[0121] Synthesized conductive polymers were analyzed with GPC (gel permeable chromatography) for measuring molecular weights thereof. The analysis was performed with GPC (Water 150 CV, column Shodex, AT-806MS (mixed column)) with NMP as solvent, 1 ml/min at 70 °C, which is recommended in NMP solvent by the manufacturer. Standard sample was polystyrene with molecular weight of 1300, 3790, 9860, 30300, 65931, 172101, 629440, and 995598.

Example 1: Preparation of Highly Conductive Polyaniline (HCPANI)

[0122] In this example, highly conductive polyaniline (HCPANI) as emeratdine base (EB) form was prepared. 100 mL of distilled and purified antine was added slowly dropwise into 6L of 1M HCl and then 4L of isopropyl alcohol was mixed with the solution. The mixed solution was maintained in the temperature of -15°C. Solution of 56 g of ammonium persulfate ((NH₄)₂S₂O₈), as an radical initiator, dissolved in 2L of 1M HCl was added slowly dropwise into the above mixed solution for 40 minutes with stirring to initiate polymerization reaction. After 3 hours, the polymerization reaction was completed to form precipitate. The obtained precipitate was filtered with filter paper and washed with 1L of 1M ammonium hydroxide (NH₄OH) solution. The precipitate was filtered with filter paper and washed with 1L of 3 mmonium hydroxide, stirred for 20 hours, washed with water, and then dried with vacuum pump for 48 hours to yield 1.5 g of polyaniline of emeraticine base (EB).

[0123] The synthesized polymer was analyzed with infrared spectroscopy and ¹³C-NMR technology. It was determined that the polyanine synthesized in this example had a peak at about 1590 cm⁻¹ of wavelength, which is assigned to the ring stretching vibration of typical quinoid ring of polyaniline, a peak at about 1495 cm⁻¹ of wavelength, which is assigned to the ring stretching vibration of typical benzenoid ring of polyaniline, and a peak at about 3010 cm⁻¹ of wavelength, which is assigned to the ring stretching vibration of C-H of aromatic ring, in infrared spectroscopy spectrum (results are not shown).

[0124] Also, it was analyzed that polyaniline had peaks at about 137 ppm and about 141 ppm of chemical shifts in ¹³C NMR spectrum, which are typical peaks of polyaniline (result not shown). Especially, polyaniline synthesized in this example had higher peak intensity at about 137 ppm than peak intensity at about 141 ppm, which is resulted from ortho- coupling, in ¹³C NMR spectrum. Therefore, it was certified that polyaniline in this example was synthesized mainly by para-coupling, not ortho-coupling and has much less side chains compared to conventional polyaniline.

55 Example 2

[0125] The procedures and conditions were repeated as example 1, except that polymerization reaction was performed at about -25 °C. Polymerization reaction were performed for 4~6 hours. It was certified that obtained material

was polyaniline of emeraldine base form through infrared spectroscopy and NMR technology (results are not shown).

Examples 3

5 [0126] The procedures and conditions were repeated as example 1, except that ammonium persulfate as a radical initiator was added drop wise for 3 hours. Polymerization reaction were performed for 3-8 hours. It was certified that obtained material was polyaniline of emeraldine base form through infrared spectroscopy and NMR technology (results are not shown).

Example 4

[0127] The procedures and condition were repeated as example 3, except that ferric chloride as a radical initiator was added to the reaction system prior to changing color of reaction vessel from blue to green. Ferric chloride was added 0.1 molar equivalents to HCI 0.1 moles. Polymerization was performed for 3~6 hours. It was certified that obtained material was polyanitine of emeraldine base form through infrared spectroscopy and NMR technology (results are not shown).

Example 5

20 [0128] The procedures and conditions were repeated as example 1, except that chloroform instead of isopropyl alcohol as the organic solvent was used. The ratios of chloroform to hydrochloric acid solution dissolving aniline monomer were respectively 2.1 and 1.1 by volume, and polymerization reaction was performed for 3-6 hours. It was certified that each of obtained materials is polyaniline of EB form with infrared spectroscopy and NMR technology (results are not shown).

Example 6

[0129] The procedures and conditions were repeated as example 1, except that mixed solvent of chloroform and isopropyl alcohol (v/v=1:1) instead of isopropyl alcohol as the organic solvent was used. The ratios of mixed solvent to hydrochloric acid solution dissolving smilline monomer were respectively 2:1 and 1:1 by volume, and polymerization reaction was performed for 3-6 hours. It was certified that each of obtained materials is polyaniline of EB form with infrared spectroscopy and NMR technology (results are not shown).

Example 7

[0130] The procedures and conditions were repeated as example 2, except that mixed solvent of chloroform and 4-methyt-2-pentanone (v/v=1:1) instead of isopropyl alcohol as the organic solvent was used. The ratio of mixed solvent to hydrochloric acid solution dissolving aniline monomer was 2:1 by volume, and polymerization reaction was performed at for 6-10 hours. It was certified that each of obtained materials is polyaniline of EB form with infrared spectroscopy and NMR technlowy (results are not shown).

Example 8

- [0131] Polyaniline substituted with alkyl group on the aromatic ring was synthesized in this example. Amine group of of-hydroxyl aniline was reacted with acetic anhydride to protect the amine group, and then protected aniline was reacted with rexame bromide in basic condition to substitute the aniline with alkyl group (frexyl group) on the aromatic ring. The protected amine group of obtained product was deprotected with hydrochloric acid to produce aniline derivative substituted with alkyl group on the aromatic ring.
 - [0132] 5 g of the synthesized aniline derivative was added drop wise into 300 mL of 1M HCI solution, and the 200 mL of dichloro methane was mixed to the solution. The solution was maintained at 5 °C, 100 mL of 1M HCI solution dissolving 1.2 g of ammonium persulfate was added drop wise to the mixed solution for 40 minutes with stirring. After 24 hours, obtained solution was separated to extract organic layer with separatory furnnel. The extracted organic layer was transferred to 200 mL of 1M ammonium hydroxide (NH₄OH), stirred for 24 hours, filtered, and then dried or 24 hours with vacuum pump to yield 1.5 g of polyaniline of emeraldine base. It was certified that obtained material is polyaniline of E8 form with infrared spectroscopy and NMR technology (results are not shown).

Example 9

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[0133] The procedure was repeated as example 1, except carboxylic acid monomer represented by structure below, which is disclosed in example 5 of international patent publication NO. WO 02-074833 as a soluble self-oriented material, was mixed to the reaction system in the ratio of 15 % by weight to aniline monomer.

[0134] It was determined that the polymer by polymerization reaction has board-shaped particles, which was different form the polymer in example 2 not adding the soluble self-oriented material. It was certified that obtained polymer was polyaniline of emeraldine base form with infrared spectroscopy and NMR technology (results are not shown).

Comparative Example 1

Preparation of Polyaniline by Conventional Method

[0135] Polyaniline of emeraldine base (EB) was prepared according to conventional MacDiarmid method (MacDiarmid et al., Conducting Polymers Ed. By Alcacer, Dordrecht, 105, 1987).

[0136] Solution of 10 mL of distilled and purified aniline dissolved in 800 mL of 1M HCl was introduced into Erlemmeyer flask. Solution of 5.6 g of ammonium persulfate dissolved in 200 mL of 1M HCl was added slowly dropwise into the flask for 15 minutes with stirring to form polyaniline. After 2 hours, the polymerization reaction was completed to obtain preopipate. The obtained precipitate was filtered with filter paper and washed with 100 mL of ammonium hydroxide. The washed precipitate was transferred to 500 mL solution of 0.1 M ammonium hydroxide, stirred for 20 hours, filtered, and dried with vacuum pump for 48 hours to yield 1.5 g of polyaniline of emeratione base. It was certified that the synthesized polymers were polyaniline as EB form with infrared spectroscopy and NMR analysis (results are not shown).

Comparative Examples 2~4: Preparation of Polyaniline

[0137] The procedures and conditions were repeated as comparative example 1, except that each of the polymerization reaction was performed respectively at -5 °C for 4 hours, -10 °C for 10 hours and at -15 °C for 17 hours with addition of 3M LiCl. It was certified that each of obtained materials is polyaniline of EB form with infrared spectroscopy and NMR technology (results are not shown).

Example 10: Measurement of Intrinsic Viscosity of Polyaniline

[0138] Highly conductive polyaniline (HCPANI) of emeraldine base form synthesized in examples 1 to 9 and conventional polyaniline (PANI) of emeraldine base from synthesized in comparative examples 1 to 4 were dedoped with ammonium hydroxide, dissolved in strong sulfuric acid of 0.1 gld. And then intrinsic viscosity (n) of HCPANI and PANI were determined at 30 °C. Table 1 shows the results of intrinsic viscosity for HCPANI and PANI. It was certified that all the synthetic compounds were polymers from measuring the intrinsic viscosity of the properties of the proper

Table 1:

Intrinsic Viscosity				
EXAMPLE	Intrinsic Viscosity (dl/g)			
1	2.2			

Table 1: (continued)

Intrinsic Viscosity					
EXAMPLE	Intrinsic Viscosity (dl/g)				
2	2.5				
3	2.4				
4	1.8				
5	2.2-2.3				
6	2.7				
7	2.9				
8	0.2				
9	1.3				
Comparative Example 1	0.8				
Comparative Example 2	1.1				
Comparative Example 3	1.1				
Comparative Example 4	1.2				

Example 11: Measurement of Optical Properties of Polyaniline

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[0139] Solid powder polyaniline HCPANI of EB form (HCPANI) synthesized in example 1 and solid powder polyaniline of EB from (PANI) synthesized in comparative example 1 were analyzed with ¹³C CPMAS-NMR and PAS spectroscopy, ¹³C CPMAS-NMR spectrum was measured at 100.6 MHz and spinning rate 7 KHz in tetramethy silane (TMS) as standard with Bruker NMR instrument. PAS spectrum was measured in helium with infrared spectrometer (Magna 550 PAS detector).

[0140] FIG. 2 shows a ¹³C CPMAS NMR analysis result of HCPANI synthesized in example 1. FIG. 3 shows a ¹³C CPMAS NMR analysis result of PANI synthesized in comparative example 1. FIG. 4 shows a PAS analysis result of HCPANI synthesized in comparative example 1. FIG. 4 shows a PAS analysis result of PANI synthesized in comparative example 1. [0141] As shown in FIG. 2, HCPANI synthesized according to example 1 has 2 remarkably distinguishable peach at around 140 ppm of chemical shift, that is, at about 138 ppm and at about 143 ppm of chemical shift is PANI should be peach at about 138 ppm of chemical shift is peach intensity at about 138 ppm of chemical shift is peach at a bout 138 ppm of chemical shift is peach at a bout 138 ppm of chemical shift is peach at a bout 138 ppm of chemical shift is peach at a bout 138 ppm of chemical shift is peach at a bout 138 ppm of chemical shift is peach at a bout 138 ppm of chemical shift is peach at a bout 138 ppm of chemical shift is peach at a bout 138 ppm of chemi

[0142] Besides, HCPANI had two relatively weak peaks at about 1107 cm⁻¹ in PAS spectrum as shown in FIG. 4, on the other hand conventional PANI had a relatively strong peak at about 1107 cm⁻¹ in PAS spectrum as shown in FIG. 5.

Example 12: Observation of Particle Formation

[0143] HCPANI powder of EB forms synthesized in above Examples 1 to 5 and PANI powder of EB from synthesized in comparative example 1 was analyzed with scanning electron microscopy in this example. FIGS. 6A, 6B, 6C, 6D, and 6E to 6F shows respectively a electron microscopy of particle structure of configuration for HCPANI powder of EB form synthesized in each of Example 1, Example 2, Example 3, Example 4, and Example 5.

[0144] As shown the figures, all of HCPANI particles synthesized according to the preferred example of the present invention show holes such as "Gomen plastics", a hollow quadra-angular rod (or har) shape like "honeycomb", or multiple layers such as "anion coat". In other words, it was analyzed that HCPANI particle of the present invention has steric configuration of much increased surface areas compared to the conventional PANI (FIGS. 7A and 7B). As shown in FIG. 6e, which is a electron microscopy enlarged by 30,000 for HCPANI particle synthesized in Example 5, particles of about 20 – 80nm were assembled to form a kind of network configuration such as bunches of grapes. In other words, it was confirmed that HCPANI produced in the present invention was synthesized as anno-sized particles to form provise network configurations. Therefore, it is expected that the conductive polymers synthesized according to the present invention is synthesized as time rachains and thereby increasing their solubility in common organic solvents.

[0145] On the other hand, polyaniline synthesized according to the convention MacDiarmid method in comparative

example 1 (PANI) has only compact structure, not network configuration, as shown in FIGS. 7A and 7B. [0146] However, HCPANI powder of EB form synthesized in Example 1 had a mean area of 145 microns and a mean volume of 230 microns, while polyamiline powder of EB form synthesized in comparative example 1 had a mean area of 7 microns and a mean volume of 18 microns, in the particle analysis by light scattering.

Example 13: Measurement of Apparent Density

[0147] In this example, the apparent density of HCPANI powder of EB forms synthesized according to above examples 1 to 6 was measured. The apparent density of polyaniline was measured by calculating injection amount of each oolvmer from specified funnel by American Society for Testion and Materials (ASTM) D1869-96.

[0148] As described in above Example 12, HCPANI has characteristic particles and pores, which increases its surface area and thereby enhancing solubility to common solvents. Also, the apparent density, which means a mass per unit volume of particle, is one important physical property.

[0149] It was measured that HCPANI synthesized according to the present invention had apparent density of 0.0495

~ 0.146 (g/cm³), which is very low compared to conventional polyaniline.

Example 14: Measurement of Electrical Conductivity of Polyaniline as Pellet

[0150] In this example, electrical conductivity of HCPANI salts synthesized in examples 1 to 9 and PANI salts synthesized in comparative examples 1 to 4 as pellets were measured as described above. It was determined that HCPANI salts synthesized in Examples 1 to 4 has electrical conductivity of 16–38 S/cm, while PANI salts synthesized in comparative examples 1 to 4 has electrical conductivity of 2–5 S/cm.

Example 15

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Measurement of Electrical Conductivity of Polyaniline in CSA Solution

[0151] In this example, both HCPANI salts synthesized in Examples 1 to 7 and PANI salts synthesized in comparative examples 1 at were dedoped to obtain polyanisine of emeration base. 1.57 g of camphor sulforinc aid (CSA) was mixed with respective 1.23 g of polyaniline EB (molar equivalent of 1:2). The mixtures are dissolved in meta-cresol with concentration of 2 % (w/w) and the solution was prepared by sonication for 2 hours. 0.5 mL of the solution was casted on side glass and offeed at 50 °C to manufacture film samples with a thickness of 0.5 ~ 80 µm. Electrical conductivity was performed on 3 film samples with a samples with a thickness of 0.5 ~ 80 µm. Electrical conductivity was performed on 3 film samples with a polyaniline film.

Table 2

Electrica	al Conductivity
EXAMPLE	Electrical Conductivity(S/cm)
1	690
2	810
3	760
4	660
5	860-920
6	1180
7	1350
8	10
9	* 510
	** 480
Comparative Example 1	210

^{*:} after polymerization reaction, soluble self oriented material is deleted.

^{**:} after polymerization reaction, soluble self oriented material is comprised.

Table 2: (continued)

Electrical Conductivity				
EXAMPLE	Electrical Conductivity(S/cm)			
Comparative Example 2	170			
Comparative Example 3	250			
Comparative Example 4	190			

Example 16: Measurement of UV-VIS-NIR Spectrum

[0152] Polyaniline of EB form synthesized in Example 2 was changed to polyaniline salt as Example 15, and then the salt was measured with UV-VIS-NIR spectrophotometer in this Example. FIG. 8 shows an analytical result of UV-VIS-NIR Spectrophotometer for emeratdine base doped with camphor sulfonic acid (EB-CSA).

[0153] In UV-VIS-NIR spectrum for polyaniline of emeraldine salt, free carrier tail which contributes to enhancing electrical conductivity of the salt usually initiates at no less than 1000 nm. However, as shown in FIG. 8, it was observed that the emeraldine salt synthesized in Example 2 does not show a localized polaron band at around 1000 nm of wavelength, but shows a continued increased absorbency line. Especially, the emeraldine salt synthesized in Example 2 had a strong polaron band by comparing the peak intensity of meta-cresol at about 300 nm to the peak intensity at about 2000 nm by polaron band.

[0154] Such a result supports the reason structurally that the emeratidine base of Example 2 had high electrical conductivity as indicated in table 2 above. The strong absorbency at near infrared rays (about 2000 nm wavelength) is similar to the absorbency for metals. In other words, it is expected that polyaniline of emeratidine salt synthesized according to the preferred examples of the present SSDP process has "true metal" properties and therefore, may be used for EMI sheld, unlike the conventional polyaniline emeration salt which has disordered properties.

Example 17

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Measurement of Molecular Weight and Distribution Degree with GPC

[0155] Polyaniline with enhanced solubility was synthesized by substituting it with LBOC in this Example according to the literature procedure (Lee et al, Macromolecules, 34, p4070, 2004). Polyaniline was prepared with the same procedures and condition as Example 1.

[0156] 1.0 g (5.5 × 10-3 mole) of the synthesized emeraldine base and 4.8 g (2.2 × 10-2 mole) of di-tert-butyloxocarbonyl (di-t-BOC) were dissolved in 30 mL of NMP. 20 mL of pyridine was added into the solution, and then the solution was stirred at 90 °C for 6 hours. The reaction product was precipitated with excessive water to filter, washed with solution of water and ethanol (1:1) to yield pure 0.6 g of t-BOC substituted polyaniline (HCPANI-tBOC). The obtained HCPANI-t-BOC was dissolved in tetrahydrofrome (THF), and then its molecular weight and molecular weight distribution are measured with got permeation chromatography (GPC, Waters C).

[0157] It was determined that t-BOC substituted polyaniline had a number average molecular weight of 44,000 and a weight average molecular weight of 46,000. Besides, it wad analyzed that t-BOC substituted polyaniline had a molecular weight distribution degree of 1.1, which means t-BOC substituted polyaniline in this example had substantially monodisperse distribution.

Example 18: NMR Analysis of Polyaniline

[0158] HCPANI synthesized according to Example 1 was analyzed by MMR in this Example. HCPANI salts of Example 1 was substituted with tert-butoxycarbony (I-BOC) to enhance solubility thereof for certifying its structures in solution state NMR analysis (C13 NMR, Jed1 YH400).

[0159] Introduction of L-BOC group into HCPANI was performed in accordance with literature (Lee et al., Macromolecules, 2004, 37, pp. 4070-4074). 4.0 g of each polyaniline powder and 13 mL of pyridine was added in 100 mL of Nrmethylpyrroidinone (NMP), into which a solution comprising 9 g of di-terl-butydicarbonate dissolved in 50 mL of NMP were added slowly at 80 °C. The mixed solution was stirred in nitrogen reflux for 3 hours to obtain product. The product was washed with methanol and dried to yield pole dark reddish powders.

[0160] Such powders were dissolved in CDCl₃, which is solvent of NMR, to obtain ¹³C NMR spectra. FIG. 9 shows an NMR analysis result of HCPANI substituted with I-BOC of the present invention. As shown the figures, it was determined that HCPANI-I-BOC had three peaks, which correspond to quaternary carbons, at about 140 ppm, about 148

ppm, and about 159 ppm of chemical shifts in solution state ¹³C NMR spectrum.

Example 19

5 Measurement of Molecular Weight and Distribution Degree with GPC

[0161] 1.0 g (5.5 × 10.3 mole) of the synthesized emeraldine base and 1.2 g of phenyl hydrazine were dissolved in 30 mL of NMP, and then reduced polyaniline was precipitated to 1L of desiccated toluene. The precipitated polyaniline was washed 3 times with toluene to yield 0.5 q of yellowish brown leucoemeraldine base powder.

[0162] The produced leucoemeraldine base was dissolved in NMP, and then its molecular weight distribution are measured with GPC (Waters Co.). Polyvinj pyridine was used to determine molecular weight and distribution degree of polyaniline as a standard sample. It was determined that leucoemeraldine base had respectively a number average molecular weight of 45,000 and 37,000, weight average molecular weight of 112,000 and 37,000, and instribution degree of 2.5 and 2.4, according to the ratio of 2.1 and 1.1 of polymer to solvent. Fig. 10 shows a distribution of molecular weight of highly conductive polyaniline synthesized according to the present invention by GPC.

[0163] Angelopoulos et al. reported that synthesizing polyaniline by conventional MacDiarmid method resulted in high molecular weight polyaniline and low molecular weight polyaniline and thereby showing various peaks. Especially, high molecular weight polyaniline comprises and/inp dut 4-10 8 among produced polymers, and synthesized polyares had broad degree of dispersion of about 3.7 to 6.6 (Angelopulos et al., Synth. Met. 84, p 35, 1997). On the other hand, the high molecular-weight polymer according to this example shows a unique peak, not separated peaks, and lower molecular distribution decree as shown in FIG. 3.

Example 20: Preparation Polypyrrole

[0164] A solution comprising 33.5 g (0.5 mol) of distilled and purified pyrrole dissolved in 1.0 L of 1 M HCl was added drop wise to 500 mL of chloroform, and then the solution was maintained in the temperature of -5 °C. Solution of 0.1 mol of ammonium persulfate ($(NH_2)_5 2_0$), as an radical initiator, dissolved in 100 mL of 1 M HCl was added slowly into the above mixed solution for 10 minutes with stirring intensively to initiate polymerization reaction. After 40 hours, the polymerization reaction was completed. The mixed solution was poured into methanol solution, and washed with distilled water several times. Residual precipitate was filtered, and then dried in vacuum oven for 24 hours. Obtained precipitate was transferred into 1. L of 1 M ammonium hydroxide, stirred 20 hours, washed with water, and then dried with vacuum pump for 48 hours to yield 11g of polypyrrole.

[0165] It was determined that polypyrrole in this example had inherent viscosity of 0.3 (in MMP) and that polypyrrole particle doped with HCl had electrical conductivity of 45 S/cm.

[0166] It will be apparent to those skilled in the art that various modifications and variations can be made in the fabrication and application of the present invention without departing from the spirit or scope of the invention. Thus, it is intended that the present invention cover the modifications and variations of this invention provided they come within the scope of the appended claims and their equivalents.

Claims

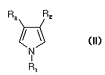
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- A process of synthesizing a conductive polymer, the process comprising:
 - (a) mixing a monomer substituted with an amine group and an organic solvent with an acid solution; and (b) adding a radical initiator dissolved in a protonic acid into the acid solution to synthesize the conductive polymer.
- The process according to claim 1, wherein the monomer substituted with the amine group is mixed with the acid solution prior to the organic solvent.
 - The process according to claim 1, wherein the monomer substituted with the amine group has a structure represented by formula I below.

wherein R₁ is hydrogen, alkyl, or alkoxy group; and each R₂ to R₃ is respectively hydrogen, alkyl, alkenyl, cycloalkyl, cycloalkenyl, alkyl-thioalkyl, alkanoyl, thioalkyl, aryl-alkyl, alkyl-amino, amino, alkoxy carbonyl, alkyl sulfonyl, alkyl sulfinyl, thioaryl, sulfonyl, carboxyl, hydroxyl, halogen, nitro, or alkyl-aryl.

 The process according to claim 1, wherein the monomer substituted with the amino group has a structure represented by formula II below.



wherein R_1 is hydrogen, alkyl, or alkoxy group; and each R_2 and R_3 is respectively hydrogen, alkyl, alkenyl, cylolalkyl, bydroalkenyl, alkyl-amino, amino, alkoxy carbonyl, alkyl-tionio, amino, alkoxy carbonyl, alkyl-tionyl, alkyl-amino, amino, alkoxy carbonyl, alkyl-tionyl, alkyl-tiony

5. The process according to claim 1, wherein the acid comprises inorganic acid.

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- The process according to claim 1, wherein the acid is selected from the group consisting of hydrochloric acid, sulfuric acid, nitric acid, or phosphoric acid.
- 7. The process according to claim 1, wherein the protonic acid comprises an inorganic acid.
- The process according to claim 7, wherein the inorganic acid is selected from the group consisting of hydrochloric acid, nitric acid, sulfuric acid, phosphoric acid, hydrofluoric acid, or hydroiodic acid or mixtures thereof.
- 9. The process according to claim 1, wherein the protonic acid comprises an organic acid.
- 10. The process according to claim 9, wherein the organic acid is selected from the group consisting of methyl sulfonic acid, dodecyl benzene sulfonic acid, antraquinone-2-sulfonic acid, 4-sulfosalicylic acid, camphor sulfonic acid, chlorinated sulfonic acid, filtifuor-sulfonic acid.
- 11. The process according to claim 1, wherein the organic solvent has a solubility parameter of about 17 to about 29.
- 12. The process according to claim 1, wherein the organic solvent comprises hydrocarbons unsubstituted or substituted with hydroxyl, halogen, oxygen, ketone, or carboxyl group.
 - 13. The process according to claim 1, wherein the organic solvent is an alkyl halide.

- 14. The process according to claim 1, wherein the organic solvent comprises dichloromethane, pentachloro ethane, 1,1,2,2-tetrachloro ethane, richloro ethane, trichloro ethylene, dichloro methane, chloroform, ethyl bromide, ethyl chloride, dichloro propane, trichloro ethane, bis(2-chloroethyl)ether, dichloro ethyl ether, 1,2-dichloro benzene, or mixtures thereof
- 15. The process according to claim 1, wherein the organic solvent is selected from the group consisting of comprise 1-propanol, 2-methyl-2-propanol, 1,2-dipropandiol, 1,3-propandiol, isopropyl alcohol, butanol, neopentanol, 2-methyl-2-butanol, 2-methyl-1-butanol, 3-methyl-1-butanol, 3-methyl-1-butanol, 3-methyl-1-butanol, 3-methyl-1-butanol, 3-methyl-1-pentanol, 3-methyl-1-pentanol, 3-methyl-1-pentanol, 3-methyl-1-pentanol, 3-methyl-1-pentanol, 3-methyl-3-pentanol, 3-methyl-3-pentanol, 3-methyl-3-pentanol, 3-methyl-3-pentanol, 4-methyl-2-pentanol, 2-methyl-3-pentanol, 3-methyl-3-pentanol, 4-methyl-2-pentanol, 2-methyl-1,3-hexandiol, octanol, 1-octanol, 2-butyl-1,3-hexandiol, octanol, 1-octanol, 2-butyl-1,3-hexandiol, octanol, 1-octanol, 2-butyl-1,3-hexandiol, octanol, 1-octanol, 2-butyl-1,3-hexandiol, octanol, 1-methyl-1-pentanolol, 1-methyl-1-pentanolol, 1-methyl-1-pentanolol, 2-butyl-1,3-hexandiol, octanol, 1-octanol, 2-butyl-1,3-hexandiol, octanolol, 1-methyl-1-pentanolol, 1-methyl-1-pentanolol, 2-methyl-1-pentanolol, 2-methyl-1-pentan

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- 16 16. The process according to claim 1, wherein the organic solvent comprises ethylene glycol monoethyl ether, ethylene glycol monomethyl ether, ethylene glycol monobutyl ether, diethylene glycol monobutyl ether of diethylene glycol monobutyl ether, diethylene glycol diethyl ether, diethylene glycol monomethyl ether, 1,4-dioxane, or mixtures thereof.
- 70 17. The process according to claim 1, wherein the organic solvent comprises butyl methyl ketone, methyl-ethyl ketone, 4-hydroxy-4-methyl-2-pentanone, cyclopentanone, diacetone alcohol, 4-methyli-pentanone, 4-methyl-2-pentanone, or mixtures thereof.
- 18. The process according to claim 1, wherein the organic solvent comprises diethyl carbonate, benzyl acetate, dimetryl glutarate, ethylacotoacetate, isobutyl isobutanate, isobutyl acetate, meta-cresol, toluene, xylene, nitrobenzene, tetrahydrofuran, N-methyl-2-povidione, dimethyl sulfoxide, NN-dimethylformamide, or mixtures thereof.
 - 19. The process according to claim 1, wherein the radical initiator comprises ammonium persulfate, hydrogen peroxide, manganese clioxide, potassium dichromate, potassium diodate, ferric chloride, potassium permanganate, potassium bromate, potassium permanganate, potassium bromate, potassium chorate. or mixtures thereof.
 - 20. The process according to claim 1, wherein step (b) is performed in the temperature of between about -45 °C to about 40 °C.
- 21. The process according to claim 1, wherein the radical initiator and the organic solvent comprises an organic phase, wherein the organic phase comprises about 5~95 % by weight based upon total aqueous solution.
 - 22. The process according to claim 2, further comprising step (c) dedoping the conductive polymer with a base.
- 23. The process according to claim 22, wherein the base comprises hydroxide compounds.
 - 24. A conductive polymer synthesized according to claim 1, wherein the conductive polymer has a hollow quadraangular rod shape and honeycombed network configuration.
- 45 25. The conductive polymer according to claim 24, wherein the conductive polymer is consisted of nanometer particles.
 - 26. The conductive polymer according to claim 24, wherein the conductive polymer is consisted of nanometer tubes.
 - 27. The conductive polymer according to claim 24, wherein the conductive polymer is consisted of nano-fibers.
 - 28. The conductive polymer according to claim 24, wherein the conductive polymer has an apparent density in the range of about 0.03 ~ 0.19 measured in ASTM Standard D1895-6.
- 29. A conductive polymer synthesized according to claim 1, wherein the conductive polymer has an electrical conductivity of at least about 300 S/cm.
 - The conductive polymer according to claim 29, wherein the conductive polymer has an electrical conductivity of at least about 500 S/cm.

- 31. The conductive polymer according to claim 29, wherein the conductive polymer has an electrical conductivity of at least about 700 S/cm
- The conductive polymer according to claim 29, wherein the conductive polymer has an electrical conductivity of at least about 900 S/cm.
- 33. The conductive polymer according to claim 29, wherein the conductive polymer has an electrical conductivity is at least about 1100 S/cm.
- 34. The conductive polymer according to claim 27, wherein the conductive polymer has an electrical conductivity of at least about 1300 S/cm.
- 35. A conductive polymer synthesized according to claim 22, wherein the conductive polymer has a hollow quadra-angular rod shape and honeycombed network configuration, wherein the conductive polymer has a repeat unit represented by the formula below and the conductive polymer has at least one single peak at about 123 ppm of chemical shift and at about 158 ppm of chemical shift in a ¹³C CPMAS NMR spectrum and/or has identifiable peaks at around 140 ppm of chemical shift in a ¹³C CPMAS NMR spectrum.

wherein x and y is respectively a molar fraction of quinonediimine structural unit and phenylenediamine structural unit in the repeating unit, and 0 < x < 1, 0 < y < 1 and x + y = 1; and n is an integer of 2 or more.

- 36. The conductive polymer according to claim 35, wherein the conductive polymer forms peaks at about 138 ppm of chemical shift and at about 143 ppm of chemical shift in a ¹³C CPMAS NMR spectrum.
- 37. The conductive polymer according to claim 35, wherein the conductive polymer has 1,36 larger than 1,43, wherein 1,38 represents a peak intensity at about 138 ppm of chemical shift in the 15C CPMAS NMR spectrum and 1,43 represents a peak intensity at about 149 ppm of chemical shift in the 15C CPMAS NMR spectrum.
 - 38. The conductive polymer according to claim 37, wherein the conductive polymer has a peak intensity ratio, I 138/ I143, of equal to or more than 1.2 in the ¹³C CPMAS NMR spectrum.
 - 39. The conductive polymer according to claim 35, wherein the conductive polymer has two peaks at about wavelength 1107 cm⁻¹ in PAS spectrum.
- 40. A polyaniline having a repeat unit represented by the formula below, wherein the polyaniline has three main peaks corresponding to quaternary carbon in a solution state ¹³C NMR spectrum in case the polyaniline is substituted with tert-butoxycarbonyl.

50 [Formula]

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wherein x and y is respectively a molar fraction of a quinonedimine structural unit and phenylenediamine structural unit in the repeating unit, and 0 < x < 1, 0 < y < 1 and x + y = 1; and n is an integer of 2 or more.

$$\begin{array}{c|c} & \begin{array}{c} 1 & 2 & 6 \\ 5 & 2 & 6 \\ 1 & 2 & 3 \\ \end{array} & \begin{array}{c} 3 & 3 \\ 3 & 3 \\ \end{array} & \begin{array}{c} 7 \\ N \\ \end{array} & \begin{array}{c} 1 \\ 2 & 1 \\ \end{array} & \begin{array}{c} 5 \\ N \\ \end{array} & \begin{array}{c} 8 & 4 \\ 4 & 4 \\ \end{array} & \begin{array}{c} 4 \\ N \\ y \end{array} & \begin{array}{c} 1 \\ n \\ \end{array} \\ \begin{array}{c} 1 \\ n \\$$

FIG. 1

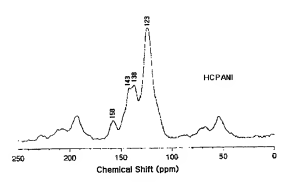
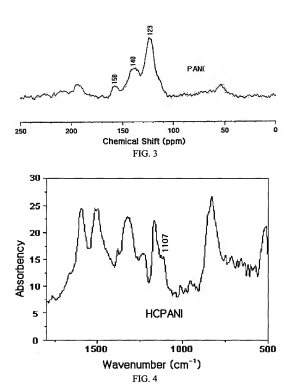
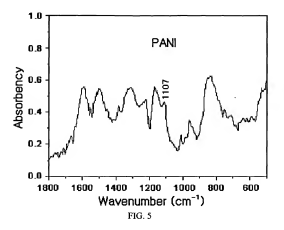


FIG. 2





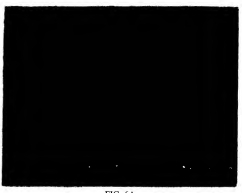
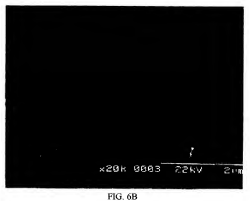


FIG. 6A



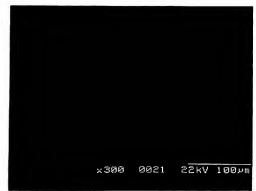


FIG. 6C

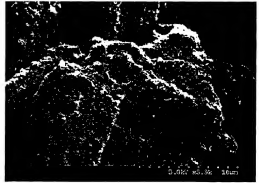


FIG. 6D



FIG. 6E



FIG. 6F

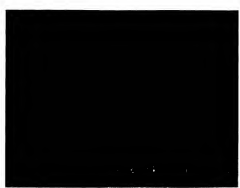
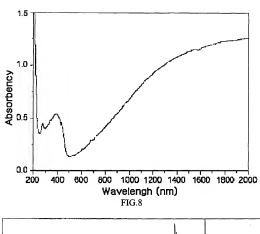
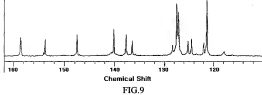


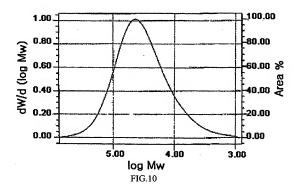
FIG.7A



FIG.7B









EUROPEAN SEARCH REPORT

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